

# Assessing metabolism and function of normothermically perfused ex vivo livers by multi-nuclear MR imaging and spectroscopy

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## Synopsis

Liver transplantation is the only cure for end-stage liver disease. Unfortunately, 20% of patients die waiting for a donor. New techniques for preserving transplant livers, such as normothermic machine perfusion (NMP), provide an opportunity to utilise 'marginal' (currently discarded) donated livers if their viability can be assessed accurately. We present initial results from a CE-marked NMP system that we adapted for use in an MRI scanner. We demonstrate the power of NMP-MRI to assess structure and metabolism in a freshly donated pig liver, dynamically over a 10-hour period. Our protocol includes <sup>1</sup>H imaging, <sup>31</sup>P spectroscopy, and hyperpolarised <sup>13</sup>C spectroscopy.

## Introduction

Liver transplantation is the only definitive treatment for end-stage liver disease, which is otherwise fatal. Waiting lists for liver transplants are increasing rapidly and there is a pressing need to use more "marginal" donated livers that are currently discarded due to the increased risk of graft failure. Recent studies have shown improved short-term post-transplant outcomes following preservation by normothermic machine perfusion (NMP).<sup>1</sup> NMP maintains the liver ex situ in an oxygenated, normothermic circuit with nutrient supplementation. However, current technology only provides crude assessment of the viability of marginal livers. To address this we combined a commercial CE-marked NMP device with a 3T MRI scanner. We show that this enables detailed metabolic and structural assessment of livers using <sup>31</sup>P-MRS, hyperpolarised <sup>13</sup>C-MRS and <sup>1</sup>H imaging techniques.

## Methods

**Perfusion system:** A CE-marked commercial NMP device ("metra", OrganOx, UK) was adapted by extending its perfusion circuit through the MRI scanner waveguide, and swapping components in the bore for non-magnetic equivalents (Fig. 1). For validation, a liver was retrieved from a 66kg white landrace pig (as per standard human donor after brain death retrieval) and cold-flushed with University of Wisconsin preservation solution before being stored on ice. The hepatic artery, portal vein, inferior vena cava and bile ducts were all cannulated. Following 4 hours of static cold storage, NMP was initiated as described by Ravikumar *et al.*<sup>2</sup> After two hours of NMP, a 15 minute period of warm ischemia was generated by disconnecting the oxygen supply. Warm ischemia was confirmed by the partial pressure of O<sub>2</sub> in the perfusate remaining below 5 kPa.

**MRI protocol:** Scans were performed on a 3T TIM-Trio (Siemens) for ten hours (Fig. 2). **<sup>31</sup>P-MRS:** a 2-channel loop-butterfly coil (Rapid Biomedical, Germany) was used to acquire non-localised FIDs repeatedly for 3.5 hours (TR/TE=3000/0.85 ms, FA=90°, bandwidth=4 kHz, points=2048, averages=200). **<sup>13</sup>C-MRS:** Nonsterile [1-<sup>13</sup>C]pyruvate was hyperpolarized with EPA radical using a 5T GE Spinlab<sup>3</sup> and injected manually into a drug line placed into the hepatic portal vein at the same time as the acquisition of slice-selective FIDs (TR=1 s, FA=10°, bandwidth=5 kHz, points=2048, slice thickness=10 cm). Spectra were acquired using a 10 cm loop coil (PulseTeq, UK). **<sup>1</sup>H-MRI:** A diffusion-weighted spin-echo sequence with echo planar readout with a 32-channel receive array (InVivo Inc, USA), and 12 b-values (0, 10, 20, 40, 60, 80, 100, 200, 300, 400, 600, 800), in 3 directions was acquired.<sup>4</sup> TR/TE=2400/65 ms, averages=2, slices=20, voxel size=1.4×1.4×8 mm<sup>3</sup>. ROIs were then drawn in the right lobe of the liver and signal intensity fitted to quantify perfusion by intravoxel incoherent motion (IVIM):<sup>5</sup>

$$\frac{S_b}{S_0} = f e^{-bD^*} + (1-f) e^{-bD},$$

where  $f$  is the perfusion fraction,  $D$  is the apparent diffusion coefficient experienced by the nonvascular compartment, and  $D^*$  is the pseudo-diffusion coefficient experienced by the microvascular compartment.

## Results

The ATP time course obtained by <sup>31</sup>P-MRS (Fig. 3) shows a 2-fold increase in the amount of ATP detected at the start of perfusion, then a gradual decrease following a 15 minute interruption to the oxygen supply. The large oxygen bolus at 2.5 hrs causes another increase in ATP. The inverse trend is seen in the inorganic phosphate (Pi). The ratios of lactate:pyruvate, alanine:pyruvate, and (bicarbonate+CO<sub>2</sub>):pyruvate following 7.5 hours of NMP were 0.104±0.004, 0.012±0.008, and 0.024±0.005 respectively. The effect of the varying production rates of metabolites can be seen in the sizes of the metabolite peaks in the spectra (Fig. 4) The IVIM results (Fig. 5) show perfusion of all liver tissue and stable perfusion over time with no significant changes between 7 and 9 hours of NMP ( $P = 0.35$ ).

## Discussion

This study introduces a new system to examine metabolic processes in *ex vivo* livers, which are active due to the *normothermic* machine perfusion. In a proof-of-principle pig liver perfusion, regeneration of ATP was seen at the beginning of the perfusion and during a large oxygen bolus following a period of warm ischemia. The relatively large amount of lactate being produced in the hyperpolarised experiment data, combined with the gradual increase in Pi during stable perfusion, suggests that this liver was partially ischemic. We are further improving the perfusion setup to obtain fully physiological normoxia. Nevertheless, IVIM maps confirm that all regions of the liver were perfused. The next step is to verify and extend our MR-protocols on discarded human livers, for which ethical approval is in place.

## Conclusions

This study introduces a new normothermic machine perfusion and MRI paradigm for studying liver metabolism *ex vivo*. This will be useful for optimising future perfusion device design, understanding the mechanisms of preservation and post-perfusion injury in human transplantation and, perhaps, in the long

term for assessing the viability of livers for transplantation.

Acknowledgements

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References

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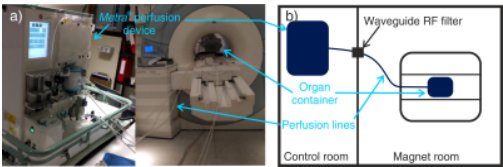
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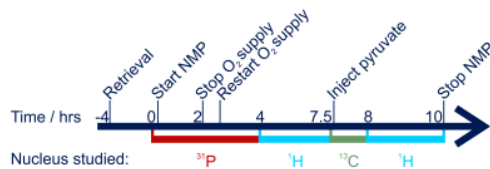
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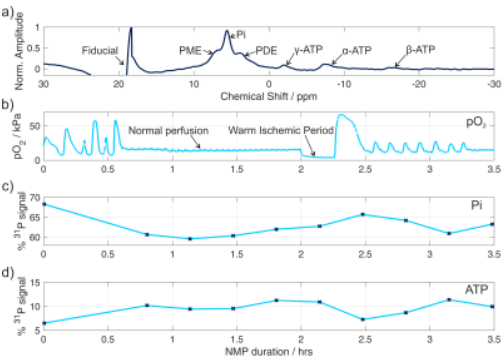
Figures



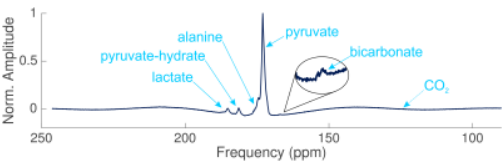
The new MR-compatible normothermic machine perfusion system based on the “metra” CE-marked commercial NMP device (OrganOx Ltd, UK).



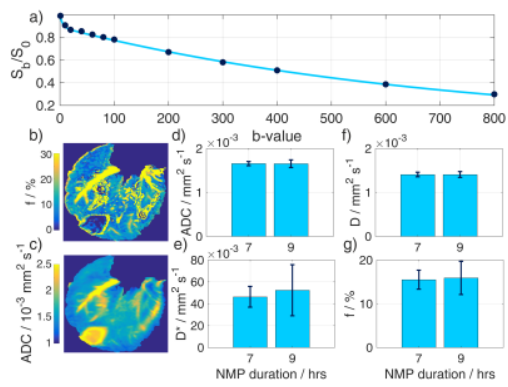
Protocol for our first perfused-liver MRI study, run in this instance on a pig liver. Longitudinal studies were performed overnight to enable 10 hours of continuous scanning.



Summary of <sup>31</sup>P-MRS results. (a) An example <sup>31</sup>P spectrum acquired following two hours of normothermic machine perfusion. (b) Variation in the partial pressure of O<sub>2</sub> in the perfusate during the perfusion. It is stable and within the physiological range once stable perfusion is established ( $pO_2 = 14.8 \pm 1.1$  kPa) then decreases to 3.7 kPa during 15 minutes of warm ischemia. (c) and (d) the variation in the signal intensity of inorganic phosphate (Pi) and adenosine triphosphate (ATP) during perfusion. Regeneration of ATP is seen at the start of perfusion and following 15 minutes of warm ischemia.



Representative spectrum from the pig liver after injection of hyperpolarised [1-<sup>13</sup>C]pyruvate. Pyruvate-hydrate, and lactate are clearly visible. Smaller amounts of alanine and bicarbonate were seen. Only a minimal amount of CO<sub>2</sub> was detected.



(a) Evolution of the signal intensity with increasing  $b$ -values showing the biexponential signal behaviour predicted by IVIM. Maps of  $f$  (b) and ADC (c) showing perfusion across all liver tissue. (d), (e), (f), and (g): Summary of IVIM data from 3 ROIs in the right lobe of the liver showing stability of the perfusion over time.